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(54) Title: METHODS OF IN SITU MODIFICATION OF PLANT GENES

(57) Abstract

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METHODS OF IN SITU MODIFICATION OF PLANT GENES

The present invention relates to the production of plants which exhibit certain desirable agronomic traits and which are produced by a non-biological process not obligatorily involving transformation or transgenesis (although these techniques can be used).

According to the present invention there is provided a method of producing plants which exhibit an agronomically desirable trait comprising mutating or otherwise modifying in situ in a plant cell at least one gene which when modified is responsible for providing the said trait and regenerating from a cell exhibiting the said trait fertile morphologically normal whole plants, characterised in that a polynucleotide is introduced into the plant cell, the said polynucleotide comprising at least one region which is substantially complementary to at least one region in the gene, which gene region when mutated or otherwise modified provides for the agronomically desirable trait, the region in the said polynucleotide containing at least one base mismatch in comparison with the like region in the said gene, so that the region in the said gene is altered by the DNA repair/replication system of the cell to include the said mismatch.

By "gene" is meant a polynucleotide comprising - contiguously - a sequence to which an RNA polymerase is capable of binding (promoter), an RNA encoding sequence and a transcription termination sequence. At least one of the following regions of the gene may be mutated or otherwise modified: promoter, RNA encoding sequence or transcription terminator. In a preferred embodiment of the method a transcription enhancing region associated with the gene is mutated or otherwise modified *in situ*.

Whilst the said trait could be an improved resistance to insects and/or fungal or bacterial infections, it is particularly preferred that the trait is herbicide resistance. The herbicides to which plants resulting from the method according to the invention are rendered resistant, or to which the said plants are tolerant or exhibit relatively improved resistance, are selected from the group consisting of paraquat; glyphosate; glufosinate; photosystem II inhibiting herbicides; dinitroanalines or other tubulin binding herbicides; herbicides which inhibit imidazole glycerol phosphate dehydratase; herbicides which inhibit acetolactate synthase; herbicides which inhibit acetyl CoA carboxylase; herbicides which inhibit protoporphyrinogen oxidase; herbicides which inhibit phytoene desaturase; herbicides which

inhibit hydroxyphenylpyruvate dioxygenase and herbicides which inhibit the biosynthesis of cellulose.

Plants which are substantially "tolerant" to a herbicide when they are subjected to it provide a dose/response curve which is shifted to the right when compared with that provided by similarly subjected non tolerant like plants. Such dose/response curves have "dose" plotted on the x-axis and "percentage kill", "herbicidal effect" etc. plotted on the y-axis. Tolerant plants will require more herbicide than non tolerant like plants in order to produce a given herbicidal effect. Plants which are substantially "resistant" to the herbicide exhibit few, if any, necrotic, lytic, chlorotic or other lesions when subjected to the herbicide at concentrations and rates which are typically employed by the agrochemical community to kill weeds in the field. Plants which are resistant to a herbicide are also tolerant of the herbicide. The terms "resistant" and "tolerant" are to be construed as "tolerant and/or resistant" within the context of the present application.

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The skilled man will appreciate that the plant material in which the *in situ* modification is performed may have been prior transformed with a gene providing for resistance to insects, fungi, and/or herbicides, or with a gene capable of providing plants regenerated from such material with, for example, an increased capacity to withstand adverse environmental conditions (improved drought and/or salt tolerance, for example) in comparison with plants regenerated from non-transformed like material.

At least one region of the polynucleotide may consist of RNA. The polynucleotide other than that comprised by the said at least one region may consist of DNA. The polynucleotide may consist of between about 30 and 250 nucleotides. In a more preferred embodiment of the polynucleotide it consists of between 50 and 200 nucleotides.

The protein encoding region of the gene may encode an enzyme selected from the group consisting of EPSPS, GOX, PAT, HPPD, ACC, ALS, BNX and protox and known mutated or variant forms thereof. In particular, the said gene may encode an EPSPS enzyme as depicted, for example, in SEQ ID Nos. 1 or 10. It is preferred that the EPSPS enzyme has least the residues Thr, Pro, Gly and Ala at positions corresponding to 174, 178, 173 and 264 with respect to the EPSPS depicted in SEQ ID No. 2, and that the said mismatch results in at least one of the following modifications in the EPSPS enzyme in comparison with the native sequence:

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- (i) Thr 174 - Ile
- Pro 178 Ser (ii)
- (iii) Gly 173 - Ala
- Ala 264 Thr (iv)

wherein (i) Thr 174 occurs within a sequence comprising contiguously Ala -Gly-Thr-Ala-5 Met; (ii) Pro 178 occurs within a sequence comprising contiguously Met-Arg-Pro-Leu-Thr; (iii) Gly 173 occurs within a sequence comprising contiguously Asn-Ala-Gly-Thr-Ala; and (iv) Ala 264 occurs within a sequence comprising contiguously Pro-Leu-Ala-Leu-Gly.

Alternatively, and/or additionally, the mismatch may result in replacement of the terminal Gly residue within the sequence motif Glu-Arg-Pro-AA1-AA2-AA3-Leu-Val-AA4-AA5-Leu-AA6-AA7-AA8-Gly- in a region of the EPSPS enzyme corresponding to that spanning positions 202 to 216 in SEQ ID No. 2 by either an Asp or Asn residue.

The plant cell to which the method of the invention is applied may be a cell of a plant selected from the group consisting of canola, sunflower, tobacco, sugar beet, cotton, maize, wheat, barley, rice, sorghum, tomato, mango, peach, apple, pear, strawberry, banana, melon, potato, carrot, lettuce, cabbage, onion, soya spp, sugar cane, pea, field beans, poplar, grape, citrus, alfalfa, rye, oats, turf and forage grasses, flax and oilseed rape, and nut producing plants insofar as they are not already specifically mentioned

The plant cell may be converted into a protoplast prior to the in situ mutation or modification of the gene - or transcriptional enhancing regions associated therewith - which when modified provides for the agronomically desirable trait.

The invention further includes plants which result from the method disclosed herein, as well as the progeny and seeds of such plants, and plant material derived from such plants. progeny and seeds.

The invention still further includes a method of selectively controlling weeds in a 25 field, the field comprising plants as disclosed in the preceding paragraph and weeds, the method comprising application to the field of a herbicide to which the said plants have been rendered resistant. Insecticidally effective amounts of insecticides and/or fungicidally effective amounts of fungicides may optionally be applied to the said plants, preferably after the herbicide has been applied to the field.

The invention will be further apparent from the following description taken in conjunction with the associated sequence listing.

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SEQ ID No. 1 shows the cDNA from Petunia encoding an EPSPS enzyme. Nucleotides 28 to 243 encode the transit peptide responsible for targeting the EPSPS enzyme encoded by nucleotides 244 to 1578 to the chloroplast. SEQ ID No. 2 shows the translational product of the sequence depicted in SEQ ID No. 1. Protein having the sequence of amino acid residues 1 to 72 constitutes the chloroplast transit peptide: protein having the sequence of amino acids 73 to 516 constitutes the EPSPS enzyme. SEQ ID Nos 3 and 4 depict peptides encoded by sequences (SEQ ID Nos 5 and 7) within exons 2 and 4 respectively of the Brassica napus EPSPS gene. Sequence ID Nos. 6 and 8 are mixed ribodeoxyribonucleic acid sequences which are capable of forming duplexes with the sequences depicted in SEQ ID Nos. 5 and 7 respectively. SEQ ID Nos. 28 and 29 are sequences which are comprised by the sequences depicted in SEQ ID Nos. 5 and 7 respectively. SEQ ID Nos. 9 and 10 depict respectively (i) the genomic DNA from Brassica napus which encodes a spliced RNA encoding an EPSPS enzyme, and (ii) the amino acid sequence of the said Brassica EPSPS enzyme. SEQ ID Nos 11 - 27 depict mixed oligonucleotides (ie containing both ribo and deoxyribonucleotides) comprising sequences (marked with asterixes in the reiteration of the sequences in the corresponding Examples) capable of causing mutations in the gene to which the oligonucleotide is targeted. The oligonucleotides depicted in SEQ ID Nos 11 to 27 are all designed to cause plant material into which they are incorporated to become resistant to herbicides, such as glyphosate and chlorsulfuron, by causing the gene encoding the proteinaceous target for the herbicide to become mutated so that the target is no longer sensitive to the herbicide. Should there by a discrepancy between the sequences depicted in the sequence listings and those corresponding sequences depicted in the Examples, the Example sequences are definitive. In the Examples sequences depicted in lower case are RNA and those in upper case are DNA.

25 Methods

Polynucleotides Mixed ribo-deoxyribonucleic acids are synthesised by synthetic and semisynthetic methods known to those skilled in the art (for example Scaringe, S.A. et al (1990), Nucleic Acids Research 18:5433-5441; Usman, N. et al (1992) Nucleic Acids Research 20:665-6699 and Swiderski, P.M. et al (1994) Anal. Biochem. 216:83-88. Eric B. Kmiec (1996) United States Patent 5,565,350). Mixed ribo-deoxyribonucleic acids are synthesised using natural nucleotides, or, in some cases, preferably with 2'-O methylated ribonucleotides. Additionally or alternatively the phosphodiester bonds of the nucleic acid

are replaced by phosphorothiodiesters or methylphosphonodiesters. Additionally or alternatively arabinose-containing nucleotides are also used.

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A duplex nucleic acid in which deoxyribonucleotides and ribonucleotides correspond with each other is termed a hybrid-duplex. When two strands form a region of duplex nucleic acid for less than all of their bases the resultant molecule is termed a heteroduplex. Two strands of a duplex can be linked by an oligonucleotide linker region to form a single polymer. The bases in the linker region are not Watson-Crick paired. A heteroduplex in which the first and second strands are portions of a single polymer is termed a hairpin duplex.

The mixed ribo-deoxyribonucleic acid useful in the present invention has at most one 3' end and one 5' end. It is constructed to contain at least one region of at least one or more usually three to four - bases that are not Watson-Crick paired. These unpaired regions form linker regions between two strands of Watson-Crick paired bases. It is preferred that the bases of the linker regions are deoxyribonucleotides.

In a preferred embodiment, the mixed ribo-deoxyribonucleic acid is constructed having two linkers arranged a) such that substantially all of the remaining bases are Watson-Crick paired and b) such that the 3' and 5' ends of the polymer are Watson-Crick paired to adjacent nucleotides of the complementary strand. These can be ligated to form a single continuous circular mixed ribo-deoxyribonucleic acid polymer.

In the present invention, the mixed ribo-deoxyribonucleic acid is used for the purpose of specifically introducing alterations (a mutation) into a target gene. The genetic site of alteration is determined by selecting a portion of the mixed ribo-deoxyribonucleic acid to have the same sequence as (to be homologous with) the sequence of the target site, hereinafter termed a homologous region. The area of differences between the sequence of the mixed ribo-deoxyribonucleic acid and the target gene is termed the heterologous region. Preferably there are two homologous regions in each mixed ribo-deoxyribonucleic acid flanking an interposed heterologous region, all three regions being present in a single continuous duplex nucleic acid. Furthermore each homologous region contains a portion of hybrid duplex nucleic acid. The portion of each hybrid-duplex is at least 4 base pairs, preferably 8 base pairs and more preferably about 20 to 30 base pairs. A dinucleotide base pair of homo-duplex may be placed within a region of hybrid duplex to allow ligation of the

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3' and 5' ends to each other. The total length of the two homologous regions is at least 20 base pairs and preferably is between 40 and 60 base pairs.

A region of homo-duplex can be disposed between the hybrid-duplex/ homologous regions of the vector. The interposed homo-duplex can contain the heterologous region. When the heterologous region is less than about 50 base pairs and preferably less than about 20 base pairs, the presence of an interposed homo-duplex is optional. When the heterologous region exceeds about 20 base pairs, an interposed homo-duplex is preferred.

The change to be introduced into the target gene is encoded by the heterologous region. The change to be introduced may be a change in one or more bases of the target gene sequence or the addition of one or more bases.

Design of polynucleotides to achieve in situ mutagenesis of EPSP synthase in Brassica napus variety Westar. It is known that the combination of mutations G101A and A192T in a Petunia EPSPS can provide for resistance to glyphosate, whilst maintaining a low Km for PEP. The equivalent residues in the sequence of the *B.napus* enzyme are (1) the second glycine occurring within the sequence LGNAGTAMRPLT (SEQ ID No. 3) where this G is amino acid 173 wherein amino acid 1 is the starting methionine of the transit peptide and (2) the third alanine occurring within the sequence MAAPLALGDVEI (SEQ ID No. 4) and consequential having the residue number 264.

The glycine residue occurs within exon 2 (part of which is shown below and is depicted as SEQ ID No. 5), the DNA coding sequence in the region being:

L G N A G T A M R P L T

ATTGAGTTGTACCTTGGGAATGCAGGAACAGCCATGCGTCCACTCACCGCTGCA

An example of the desired mutation is GGA ---> GCA

25 example, on one of its strands, a sequence comprising mainly of RNA which is complementary to all or part of the above DNA sequence. This RNA is interposed by a short region of DNA also complementary with the corresponding region of the above DNA sequence except for the inclusion of the specific mismatch of having a guanosine base opposite the guanosine base within the target GGA codon. A suitable mixed ribodeoxyribonucleic acid could thus include all or part of the following sequence (depicted as SEQ ID No. 6 in the sequence listing). Note that RNA sequence is marked in bold.

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TTGTACCTTGGGAATGCAGGAACAGCCATGCGTCCACTC

AACAUGGAACCCUUACGTCGTTGUCGGUACGCAGGUGAG

The corresponding alanine residue occurs within exon 4 (part of which is shown below and is depicted as SEQ ID No. 7).

M A A P L A L G D V E I

ACTGCCCTCCTCATGGCAGCTCCTTTAGCTCTTGGAGACGTGGAGATTGAGATCATT

An example of the desired mutation is GCT ---> ACT. The mixed ribodeoxyribonucleic acid designed to elicit this change includes, for example, on one of its strands, a sequence comprising mainly of RNA which is complementary to all or part of the above DNA sequence. This RNA is interposed by a short region of DNA also complementary with the corresponding region of the above DNA sequence except for the inclusion of the specific mismatch of having a thymine base opposite the guanosine base within the target GCT codon. The desired polynucleotide thus includes all or part of the RNA sequence depicted below and in SEQ ID No. 8. Note that RNA sequence is marked in bold.

TCCTCATGGCAGCTCCTTTAGCTCTTGGAGACGTGGAGATT

AGGAGUACCGUCGAGGAAATTGAGAACCUCUGCACCUCUAA

Besides the examples detailed above there will of course be many other specific changes which could be introduced into those sequences which regulate gene expression and for which polynucleotides can easily be designed by methods directly analogous to that described above and which, for example, could be useful to achieve increased expression of EPSPS. The skilled man will appreciate that many methods could be used to specify those changes potentially useful for increasing the expression of EPSPS. For example:

(1) The skilled man will be aware of instances of resistance to glyphosate having occurred in both field populations of weeds (e.g Australian Iolium) and upon continuous selection of cultured plant cells (e.g. Hollander-Czytko et al (1988) in Plant Mol. Biol, 11, 215-220; Hollander-Czytko et al (1992) Plant. Mol. Biol. 20, 1029-1036) or, for example, cultivars of birdsfoot trefoil (Boerboom et al (1990) Weed. Sci., 38, 463-467) upon glyphosate. In the latter two cases selection was shown to have resulted in a significant increase in expression of EPSP synthase. In the example of the work on cell cultures of Corydalis sempervirens (Hollander-Czytko et al (1988) in Plant Mol. Biol, 11, 215-220) a 30-40 fold increase in the cellular content of EPSP synthase and an 8-12 fold increase in transcript levels was observed. There was no amplification of the EPSP synthase gene.

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It is a routine matter in all of the above examples using methods known to the skilled—man to isolate cDNA encoding the EPSP synthases, to use these cDNA's as probes to identify clones from genomic libraries and to sequence the corresponding EPSP synthase genes and their 5' upstream and 3' downstream regions. Alternatively, genomic sequences may be isolated directly using heterologous probes and/or combinations of degenerate and inverse PCR. By comparing the sequences so obtained from 'high EPSP synthase expression' lines of plants, cultivars or plant cells with the appropriate unselected controls the specific mutation(s) responsible for conferring high expression of EPSP synthase will be identified.

- (2) Another example of a suitable method for identifying mutations potentially useful for increasing the expression of EPSP synthase is to directly select various lines of cultured plant cells or protoplasts from plant species of interest (e.g. *Brassica napus*) on increasing concentrations of glyphosate. This can be done with or without the addition of a suitable chemical mutagen. Glyphosate-tolerant lines so obtained are analysed for expression of EPSP synthase, for the level of translatable EPSP synthase gene transcript (e.g by Northern analysis) and for possible amplification of the EPSPS gene (e.g. by Southern and dot blot analysis). Cell lines of particular interest would be those where EPSP synthase was overexpressed and where this increase could not be accounted for through gene amplification. Identification of the specific mutation(s) responsible for conferring high expression of EPSP synthase are then identified as described in (1) above.
- (3) A further example of a method useful to specify mutations causing high expression of EPSPS comprises (a) subcloning the plant EPSP synthase promoter, 5' upstream sequence region, translational start region and sequence encoding the N-terminus region of EPSP synthase into a translational fusion construct directing the synthesis of a suitable and easily measurable reporter gene such as (Beta glucuronidase) (b) further cloning this into a shuttle vector containing an origin for replication in *E. coli* and also designed for site specific integration into the yeast genome (YIP), or the genome of any other suitable test cell, such that integration into a specific location can be positively selected, by for example, complementation of an auxotrophic mutation. A library of many variants specifically within the promoter and 5' upstream region of the so-designed shuttle vector is then created by mutagenesis through, for example, Mn2+-poisoned PCR of the region and maintained in *E. coli*. Members of the library are then tested by transformation into yeast. The best

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expressers in yeast are identified by increased expression of the reporter gene. The integrated DNA from these high expresser lines is then extracted, sequenced and compared with the original sequence in order to identify those specific mutation(s) which conferred increased expression. Such mutations may affect conserved domains within the promoter which bind the transcriptional activators required for gene expression. Studies of this sort may teach those skilled in the art to modify the equivalent conserved regions in other crop plant species, thus enabling the technology to be applied broadly.

The polynucleotides comprising the RNA sequences disclosed above are transfected into protoplasts of *Brassica napus* which are then cultured and subjected to the herbicide glyphosate at concentrations which are sufficient to kill like protoplasts which have not been transfected and like protoplasts which have been transfected but with a polynucleotide not comprising regions designed to elicit a mutation in the *Brassica* genome. Those transfected protoplasts which survive the herbicide at concentrations which kill the control protoplasts are regenerated into plants using known means. The increased resistance to the herbicide of the thus regenerated plants is inherited in a Mendelian manner amongst the progeny of these plants.

The skilled man will appreciate that the invention is not limited to that specifically described above in respect of the production of glyphosate resistant *Brassica napus*. For plant species for which the EPSP synthase gene sequence(s) are already available on public databases the RNA and DNA elements of the polynucleotides can easily be designed by a method directly analogous to that described for *B. napus*. Polynucleotides comprising these RNA and DNA elements can then be introduced into regeneratable plant material from other species. Moreover, the skilled man is capable of designing:

(i) polynucleotides for the in situ mutagenesis of the DNA bases flanking the translational start site to improve post transcriptional efficiency of expression of EPSP synthase in plants, for example Brassica napus variety Westar. The consensus sequences for the regions immediately surrounding the translational start sites in animals (M Kozak, 1986, Cell, 44, 283-292) and plants (G Heidecker and J Messing, 1986, Ann. Rev. Plant Physiol., 37, 439-466; V Pautot et al., 1989, Gene, 77, 133-140) have been described. It is therefore possible that improved levels of expression of the native B. napus EPSP synthase gene may be improved in situ by designing mixed ribo-deoxyribonucleic oligonucleotides to make the

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desired mutational changes, at positions -3 and + 6 as shown below. Note that conserved consensus sequences are underlined.

B. napus
$$\underline{A}$$
 \underline{A} \underline{C} \underline{A} \underline{A} \underline{T} \underline{G} \underline{G} \underline{C} \underline{G} \underline{C} \underline{C} Concensus \underline{A} \underline{A} \underline{C} \underline{A} \underline{A} \underline{T} \underline{G} \underline{G} \underline{C} \underline{C} \underline{C}

It will be obvious to those skilled in the art that this approach need not be confined to the EPSP synthase gene from B. napus, but may be applied to any plant species in which an increase in expression of the target gene is sought.

ii) polynucleotides for the *in situ* mutagenesis of the DNA bases to achieve an increase in transcriptional efficiency of expression of EPSP synthase. An approach similar to that described above may be adopted to achieve an enhancement in the rate of transcription of EPSP synthase genes by mutating bases at the "TATA" box region upstream from the transcription start point, and at the transcription start point itself. Identification of the transcription start point is identified using techniques, such as primer extension analysis, known to those skilled in the art. The "TATA" box is generally found 16-54 bases upstream of the transcriptional start. Consensus sequences have been published for plant transcription start point (V Pautot et al., 1989, Gene, 77, 133-140)

Plant Consensus CTCATCA

and "TATA" box regions (V Pautot et al., 1989, Gene, 77, 133-140)

Plant Consensus TCACTATATATAG

In both cases highly conserved bases are underlined. Comparisons between the consensus and native sequences of target EPSP synthase genes will enable bases suitable for mutational change to be identified.

(iii) polynucleotides for in situ mutagenesis to alter expression of EPSP synthase in plants, for example Brassica napus variety Westar.

Such designed polynucleotides can be introduced into totipotent plant material by known means which is then regenerated into plants which are subjected to a selection procedure to isolate those that exhibit the desired trait.

The skilled man will appreciate that directly analogous methods to those described above for EPSP synthase and glyphosate could be applied to other combinations of selecting herbicide and target gene where the aim is to specify mutations conferring over-expression.

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The invention will be further apparent from the following Examples. Throughout the Examples the expression "selecting concentrations" of herbicide is present. By this is meant a concentration of herbicide which is sufficient to kill non-transformed material, or material which otherwise does not contain the oligonucleotides which are contained within like experimental material. The skilled man will know what those concentrations are having regard to the specific circumstances relating to his particular germplasm, transformation protocols and the expected variation between replicate procedures. The oligonucleotides shown below (SEQ ID Nos 11 to 27) are all synthesised according to Yoon *et al.* (1996). In each of the Examples where the constructs contain bases depicted in lower case, the sequence comprising such bases is to be understood as being RNA, and sequences comprising bases depicted in upper case as being DNA.

Example 1 This Example demonstrates the production of corn (maize) which is resistant to the herbicide chlorsulfuron.

TGCGCG gauacuagggATTACcaccccgaaT
T
T
T
TCGCGC CTATGATCCCTAATGGTGGGGCTTT
20 3'5'

The above oligonucleotide (SEQ ID No. 11) conveniently may be introduced into corn using silicon carbide whiskers, pollen harbouring the oligonucleotide or *via* pollen tubes.

Whiskers The so called whiskers technique is performed essentially as described by Frame et al., (1994 Plant J. 6 941 -948). The oligonucleotide (1-100 µg) depicted in SEQ ID No.11 is added to the whiskers and used to transform A188 x b73 cell suspensions. The oligonucleotide(s) may be co-incubated with plasmids comprising sequences encoding proteins capable of forming recombinase complexes in plant cells such that recombination is catalysed between the oligonucleotide and the target sequence. Plant regeneration is performed using selective concentrations of chlorsulfuron in place of bialophos. Plants are transferred to pots and matured in the green house. Kernals from these plants are germinated in soil and sprayed with a selecting concentration of chlorsulfuron 9 to 14 days post emergence.

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Pollen transformation Maize pollen is bombarded with gold particles by techniques known to the skilled man. Gold particles are coated with the oligonucleotide depicted in SEQ ID No. 11. The oligonucleotides may be co-incubated with plasmids comprising sequences encoding proteins capable of forming recombinase complexes in plant cells such that recombination is catalysed between the oligonucleotide and the target sequence. Suitable bombardment methods vary in precise detail but the basic procedure is well known to the skilled man and it is thus not necessary to describe it here. Bombarded pollen is applied to receptive silks of detassled plants. Sufficient replicas are performed to pollinate a large number of plants (typically up to 500). Progeny of the plants are screened for chlorsulfuron resistant members of the population by spraying with selecting concentrations of chlorsulfuron.

Pollen tube mediated transformation Emasculated corn plants are used. Wild type pollen is applied to pollination receptive silks. After between 30 min to 6 hours the silks are cut to within one cm of the base. The above SEQ ID No. 11 oligonucleotide (1-100 µg/ 10 µl in TE) is applied to the cut surface using a 1 ml syringe and needle such that the surface is completely covered. The oligonucleotides may be co-incubated with plasmids comprising sequences encoding proteins capable of forming recombinase complexes in plant cells such that recombination is catalysed between the oligonucleotide and the target sequence. The plants are then grown in a green house with an initial humidity of about 75 %. Progeny of the plants are screened for chlorsulfuron resistant members of the population by spraying with selecting concentrations of the herbicide.

Plants derived from material into which the oligonucleotides have been incorporated are resistant, more resistant or tolerant to the herbicide, when compared to plants derived from material not containing the said oligonucleotide.

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Example 2 This Example demonstrates the production of *Arabidopsis thaliana* which is resistant to the herbicide glyphosate (and suitable salts thereof). The following oligonucleotides (depicted as SEQ ID Nos 12 to 16 in the sequence listing) are prepared using standard technology.

T to I

```
T GCGCG cauuacguccTTATCguuacgcagg T
  5
                                           Ť
      T CGCGC GTAATGCAGGAATAGCAATGCGTCC T
            3'5'
                     (SEQ ID No. 12)
     T to I2
 10
      T GCGCG cauuacgtccTTATCguuacgcaag T
     T CGCGC GTAATGCAGGAATAGCAATGCGTTC T
15
                     (SEQ ID No. 13)
    P to S
20
     T GCGCG ugucguuacgCAAGTgaauggcgac T
    Т
     T CGCGC ACAGCAATGCGTTCACTTACCGCTG T
25
            3'5'
                    (SEQ ID No. 14)
    P to S 2
      GCGCG uaucguuacgCAAGTgaauggcgac T
30
    Т
      CGCGC ATAGCAATGCGTTCACTTACCGCTG T
           3'5'
                    (SEQ ID No. 15)
35
     T GCGCG cauuacguccTTATCguuacgCAAGTgaguggcgac T
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    T
```

These oligonucleotides are introduced into *Arabidopsis* by microprojectile bombardment or protoplast uptake.

T CGCGC GTAATGCAGGAATAGCAATGCGTTCACCGCTG T 3'5' (SEQ ID No. 16)

Bombardments Arabidopsis is transformed essentially using a modified procedure as described by Seki et al. ((1991) Appl. Microbiol. Biotechnol. 36 228-230). Arabidopsis thaliana genotype C24 seeds are surface sterilised and sown on B-5 medium

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(Gamborg et al., 1968) solidified with 0.6 % agarose. The plants are grown aseptically for 4-- 6 weeks under 16 h light 8 h dark at 26 °C. Roots are harvested and cut into sections that are 0.5 - 1.0 cm long and placed onto a filter paper on medium containing B5 salts and vitamins, 3 % sucrose, 0.5 mg/ml 2,4-dichloropheonoxyacetic acid, 0.05 mg/l kinetin and 0.8 % agarose (0.5 - 0.05 medium). After two to five days the roots are ready for bombardment. Gold particles (10 mg; Hereus, 0.4-1.2 um diameter) are coated with 1 - 100 µg of oligonucleotide as follows. The oligonucleotides may be co-incubated with plasmids comprising sequences encoding proteins capable of forming recombinase complexes in plant cells such that recombination is catalysed between the oligonucleotide and the target sequence. The particles are suspended in 1 ml of absolute ethanol and incubated for three hours at room temperature then stored at -20oc. Twenty to thirty-five μl of sterile resuspended particles are collected by centrifugation in a microcentrifuge. The particles are washed with one ml of sterile distilled water and re-collected by centrifugation. Microprojectiles are then resuspended in 30 μ l oligonucleotide solution (1 -100 μ g), 25 μ l of 1M CaCl2 is added followed by 10 μl of 0.1 M spermidine (free base). The mixture is incubated on ice for 10 minutes. 1-10 µl of this solution is used per bombardment. A suitable mixture or combination of oligonucleotides is introduced into plant material either simultaneously or sequentially. If the oligonucleotides are introduced sequentially, they must be introduced in such a way that the mutation governed by the first oligonucleotide is not negated by the mutation governed by a subsequently introduced oligonucleotide. For example, if the oligonucleotide depicted by SEQ ID No. 12 is introduced first, the oligonucleotide depicted by SEQ ID No. 15 should be used subsequently. Alternatively, a single oligonucleotide comprising regions providing for multiple mutations (such as that depicted in SEQ ID No. 16) may be used.

The roots are bombarded with oligonucleotide-coated particles by a helium-driven biolistics PDS 1000 system (BioRad) with a 300 mm Hg vacuum. The levels between the rupture disk and the macrocarrier and the macro-carrier and sample are varied for maximal transformation efficiency. Rupture disks of between 1000 and 2000 psi are used. Two suitable oligonucleotides are introduced into *Arabidopsis* plant material either simultaneously or sequentially. For simultaneous transformation the oligonucleotides are used in equal molar concentrations and may be introduced into the material by multiple firings into the same tissue. For sequential transformation the roots receive at least one

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bombardment with each oligonucleotide but multiple firings of each oligonucleotide are usedif necessary to optimise transformation efficiencies.

After the bombardments the plant material is transferred to 0.5 - 0.05 medium and incubated at 26oc for one to 5 days. Regeneration of transformed material into *Arabidopsis* plants is performed as Seki *et al* 1991 with the exception that kanamycin or gentamycin are not included in any of the media. Instead the transformed material is selected by its resistance or tolerance to glyphosate, present in the selection medium at a concentration sufficient to kill control material which has been subjected to a like transformation procedure with the *proviso* that it does not contain the oligonucleotides specified above.

DNA uptake by protoplasts incubated in PEG — The protocol of Dam et al. (1989 Mol Gen. Genet 217 6-12) is followed. Instead of using linearised plasmid DNA in the transformation an equal molar ratio mix of the two oligonucleotides (SEQ ID Nos 12 and 15) are used (1-100 µg) with 50-100 µg calf thymus carrier DNA. The oligonucleotides may be co-incubated with plasmids comprising sequences encoding proteins capable of forming recombinase complexes in plant cells such that recombination is catalysed between the oligonucleotide and the target sequence. Glyphosate selection instead of hygromycin selection is applied at the same stage during callus formation. The concentration of glyphosate used is varied to give optimum selection of transformed *Arabidopsis* plants, but is determined by reference to suitable control experiments.

Plants derived from material into which the oligonucleotides have been incorporated are resistant, more resistant or tolerant to the herbicide, when compared to plants derived from material not containing the said oligonucleotide.

Example 3 This Example demonstrates the provision of glyphosate resistant Brassica napus

T to I

T GCGCG ccuuacguccTTATCgcuacgcagg T
T
T
T T CGCGC GGAATGCAGGAATAGCCATGCGTCC T
3'5' (SEQ ID No. 17)

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T to I2
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```
T GCGCG ccuuacgtccTTATCgcuacgcaag T
   T
5
   Т
                                          Т
    T CGCGC GGAATGCAGGAATAGCCATGCGTTC T
           3'5'
                   (SEQ ID No. 18)
```

10 P to S

```
GCGCG ugucgguacgCAAGTgaguggcgac T
    T
     T CGCGC ACAGCCATGCGTTCACTCACCGCTG T
15
                    (SEQ ID No. 19)
```

P to S 2

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20 T GCGCG uaucgguacgCAAGTgaguggcgac T T CGCGC ATAGCCATGCGTTCACTCACCGCTG T 25 (SEQ ID No. 20)

T GCGCG ccuuacguccTTATCgcuacgCAAGTgaguggcgac T 30 Т Т Т T CGCGC GGAATGCAGGAATAGCCATGCGTTCACCGCTG T (SEQ ID No. 21)

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These oligonucleotides are designed to target the Brassica napus EPSPS gene. The oligonucleotides provide for two changes in the sequence of the protein encoded by the gene, viz. at T102 and P106 of the Brassica mature enzyme such that the mutant gene (via an altered protein product) confers resistance to glyphosate.

The oligonucleotides are introduced into Brassica napus using known methods which 40 includes microprojectile bombardment or uptake of DNA by protoplasts.

Seeds of B.napus cv Westar are surface sterilised in 1% sodium Bombardments hypochlorite for 20 minutes. The seeds are then washed in sterile water three times and planted at a density of about 10 seeds per plate on Murashige Skoog (MS) minimal organics medium (GibcoBrl) with 3% sucrose and 0.7% phytagar (Gibco) pH 5.8. Seeds are germinated at 24 °C in 16 h light/8h dark. After five days the cotyledons are excised in such a

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way that they include approximately 2 mm of petiole at the base. Care is taken to exclude the apical meristem. The excised cotyledons are placed on MS medium, 3 % sucrose and 0.7 % phytagar enriched with 20 μ M bezyladenine with the petioles imbedded to a depth of 2 mm in the medium at a density of about ten cotyledons per plate.

Gold particles (10 mg; Hereus, 0.4-1.2 um diameter) are coated with 1 - 100 µg of oligonucleotide (SEQ ID No. 22 for example, or SEQ ID Nos. 18 and 20) in plant cells. The oligonucleotides may be co-incubated with plasmids comprising sequences encoding proteins capable of forming recombinase complexes in plant cells such that recombination is catalysed between the oligonucleotide and the target sequence. The particles are suspended in 1 ml of absolute ethanol and incubated for three hours at room temperature then stored at 20oc. Twenty to thirty five µl of sterile resuspended particles are collected by centrifugation in a microcentrifuge. The particles are washed with one ml of sterile distilled water and recollected by centrifugation. Microprojectiles are then resuspended in 30 µl solution (containing oligonucleotides depicted in SEQ ID Nos. 18 and 20, for example in an amount of about 1 -100 µg). 25 µl of 1M CaCl2 is added followed by 10 µl of 0.1 M spermidine (free base). The mixture is incubated on ice for 10 minutes. 1 -10 µl of this solution is used per bombardment.

The cotyledons are bombarded with oligonucleotide-coated particles by a helium-driven biolistics PDS 1000 system (BioRad) with a 300 mm Hg vacuum. The levels between the rupture disk and the macrocarrier and the macro-carrier and sample are varied for maximal transformation efficiency. Rupture disks of between 1000 and 2000 psi are used. The two oligonucleotides are introduced into the *Brassica* plant material either simultaneously or sequentially. For simultaneous transformation the oligonucleotides are used in equal molar concentrations and may be introduced into the explant by multiple firings into the same tissue. For sequential transformation the explants receive at least one bombardment with each oligonucleotide but multiple firings of each oligonucleotide are used as necessary to optimise transformation efficiencies.

After bombardment the explants are placed onto regeneration medium comprising MS medium supplemented with 20 μ M benzyladenine, 3% sucrose 0.7% phytagar pH 5.8. After 2 - 5 days the cotyledons are transferred to plates containing the same media but including selective concentrations of glyphosate. The petioles remain embedded in the media. The explants are left for 2 - 6 weeks and then transferred onto MS medium

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supplemented with 3 % sucrose, 0.7% phytagar pH 5.8 and selecting concentrations of glyphosate. One to three weeks later surviving shoots are transferred to rooting media which comprises MS medium, 3% sucrose, 2 mg/ml indole butyric acid, 0.7% phytagar with no glyphosate. Once roots are visible the plants are transferred to pots and propagated in the greenhouse.

Protoplast uptake The method of Golz et al. ((1990) Plant Mol Biol 15 475 - 483) is followed. Brassica napus genotype H1 is used. Instead of using plasmid DNA in the transformation an equal molar ratio mix of the two oligonucleotides (SEQ ID Nos 18 and 20) are used (1-100 µg) and 20-100 µg calf thymus carrier DNA. The oligonucleotides may be co-incubated with plasmids comprising sequences encoding proteins capable of forming recombinase complexes in plant cells such that recombination is catalysed between the oligonucleotide and the target sequence. Glyphosate selection instead of hygromycin selection is applied at the same stage during callus formation. The concentration of glyphosate used is varied to give optimum selection of transformed Brassica plants.

Plants derived from material into which the oligonucleotides have been incorporated are resistant, more resistant or tolerant to the herbicide, when compared to plants derived from material not containing the said oligonucleotide.

Example 4 This Example demonstrates the provision of corn resistant to the herbicide glyphosate (and salts thereof).

T to I

T GCGCG ccuuacgaccTTAGCGuuacgccggua T
T
T
T
T CGCGC GGAATGCTGGAATCGCAATGCGGCCAT T
3'5' (SEQ ID No. 22)

**
T GCGCG ccuuacgaccTTAGCGuuacgccagua T
T
T
T
T
T
T CGCGC GGAATGCTGGAATCGCAATGCGGTCAT T
35' (SEQ ID No. 23)

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```
P to S
       GCGCG gacquuacgCCAGTaacugucqucq T
    Т
                                           Т
5
     T CGCGC CTGCAATGCGGTCATTGACAGCAGC T
                    (SEQ ID No. 24)
    P to S 2
10
      GCGCG agcguuacgCCAGTaacugtcgucg T
                                          Т
    T
                                          Т
     T CGCGC TCGCAATGCGGTCATTGACAGCAGC T
15
           3/5/
                    (SEQ ID No. 25)
```

T GCGCG ccuuacgaccTTAGCGuuacgCCAGTaacuqucqucq T 20 Т Т T CGCGC GGAATGCTGGAATCGCAATGCGGTCATTGACAGCAGC T 3151 (SEQ ID No. 26)

These oligonucleotides which are designated as SEQ ID Nos 22-26 in the sequence listing 25 and which are produced by means known to the skilled man, may be introduced into corn using silicon carbide whiskers, pollen harbouring oligonucleotides or via pollen tubes. This transformation is performed essentially as described by Silicon carbide whiskers Frame et al. (1994 Plant J. 6 941-948). The oligonucleotide depicted as SEQ ID No 26 (1-100 µg) is added to the whiskers and used to transform A188 x B73 cell suspensions. The 30 oligonucleotides may be co-incubated with plasmids comprising sequences encoding proteins capable of forming recombinase complexes in plant cells such that recombination is catalysed between the oligonucleotide and the target sequence. Plant regeneration is performed using selective concentrations of glyphosate in place of bialophos. Plants are transferred to pots and are then matured in the green house. Caryopsis from these plants are 35 germinated in soil and sprayed with a selecting concentration of glyphosate 9 to 14 days post emergence.

Maize pollen is bombarded with gold particles (essentially as Pollen transformation. described in the above Examples) coated with a mixture of the above oligonucleotides (SEQ ID Nos 23 and 25). The oligonucleotides may be co-incubated with plasmids comprising sequences encoding proteins capable of forming recombinase complexes in plant cells such

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that recombination is catalysed between the oligonucleotide and the target sequence. Bombarded pollen is applied to receptive silks of detassled plants. Sufficient replicas are performed to pollinate a large number (typically up to 300) of plants. Progeny of the plants are screened for glyphosate resistant members of the population by spraying with selecting concentrations of glyphosate.

Pollen tube mediated transformation Emasculated corn plants are used. Wild type pollen is applied to pollination receptive silks. After between 30 min to 6 hours the silks are cut to within one cm of the base. Suitable mixtures of the above oligonucleotides (1-100µg/10 µl in TE) are applied to the cut surface using a 1 ml syringe and needle such that surface is completely covered. The oligonucleotides may be co-incubated with plasmids comprising sequences encoding proteins capable of forming recombinase complexes in plant cells such that recombination is catalysed between the oligonucleotide and the target sequence. The plants are then grown in a green house with an initial humidity of about 75 %. Progeny of the plants are screened for glyphosate resistant members of the population by spraying with selecting concentrations of glyphosate.

Plants derived from material into which the oligonucleotides have been incorporated are resistant, more resistant or tolerant to the herbicide, when compared to plants derived from material not containing the said oligonucleotide.

Example 5 This Example demonstrates the provision of tomato plants resistant to a bleaching herbicide designated as R390244.

This oligonucleotide (SEQ ID No. 27) is designed to target the codon for arginine 307 of the tomato phytoene desaturase (PDS) gene and introduce a mutation such that the mutant PDS is resistant to the herbicide R390244. The oligonucleotides may be co-incubated with plasmids comprising sequences encoding proteins capable of forming recombinase complexes in plant cells such that recombination is catalysed between the oligonucleotide and the target sequence. The oligonucleotide is introduced into tomato Mill cv H722 via microprojectile bombardment essentially as described by Eck et al. (1995 Plant Cell Reports

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14, 299-304) and as outlined above for the other crops subjected to this transformation procedure.

Regenerable cotyledon explant material (as described by Fillati *et al.* (1997). Bio/technology 5 726-730) suspensions are bombarded with SEQ ID No. C oligonucleotide-coated particles by a helium-driven biolistics PDS 1000 system (BioRad) with a 300 mm Hg vacuum. The levels between the rupture disk and the macrocarrier and the macro-carrier and sample are varied for maximal transformation efficiency. Rupture disks of between 1000 and 2000 psi are used. The oligonucleotide may be introduced into the explant by multiple firings into the same tissue as necessary to optimise transformation efficiencies. The regenerable cotyledons are bombarded at the same stage as when *Agrobacterium* is used in the method of Beaudoin and Rothstein (1997 Plant Mol Biol 33 835 -846). Regeneration of tomato plants is as described by Beaudoin and Rothstein except that no selection agent is used. Primary putative transformants are grown in the greenhouse and cuttings are propagated in soil. These cuttings, once established, are sprayed with selecting concentrations of R390244 and allow transformed herbicide resistant plants to be identified. These transformed plants are grown to maturity and seeds resulting from self pollination are collected.

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Mutation events in individuals is confirmed by amplifying the particular mutant gene sequence from herbicide resistant individuals spanning the region of mutation by PCR and sequencing individually isolated and cloned sequences.

Plants derived from material into which the oligonucleotides have been incorporated are resistant, more resistant or tolerant to the herbicide, when compared to plants derived from material not containing the said oligonucleotide.

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SEQUENCE LISTING

	(1) GENERAL INFORMATION:	
5	(i) APPLICANT: (A) NAME: ZENECA LTD (B) STREET: 15 Stanhope Gate (C) CITY: LONDON (E) COUNTRY: GB (F) POSTAL CODE (ZIP): W1Y 6LN	-
	(ii) TITLE OF INVENTION: IMPROVEMENTS IN OR RELATING TO ORGANIC COMP	OUNDS
15	(iii) NUMBER OF SEQUENCES: 29	
20	<pre>(iv) COMPUTER READABLE FORM: (A) MEDIUM TYPE: Floppy disk (B) COMPUTER: IBM PC compatible (C) OPERATING SYSTEM: PC-DOS/MS-DOS (D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)</pre>	
	(2) INFORMATION FOR SEQ ID NO: 1:	
25	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1944 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown 	
30	(ii) MOLECULE TYPE: cDNA	
	(iii) HYPOTHETICAL: NO	
35	(iv) ANTI-SENSE: NO	
	<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: Petunia hybrida</pre>	
40	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION:281578	
45	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:	
	GAATTCCCTC AATCTTTACT TTCAAGA ATG GCA CAA ATT AAC AAC ATG GCT Met Ala Gln Ile Asn Asn Met Ala 1 5	51
50	CAA GGG ATA CAA ACC CTT AAT CCC AAT TCC AAT TTC CAT AAA CCC CAA Gln Gly Ile Gln Thr Leu Asn Pro Asn Ser Asn Phe His Lys Pro Gln 10 15 20	99
55	GTT CCT AAA TCT TCA AGT TTT CTT GTT TTT GGA TCT AAA AAA CTG AAA Val Pro Lys Ser Ser Ser Phe Leu Val Phe Gly Ser Lys Leu Lys 30 35 40	147
60	AAT TCA GCA AAT TCT ATG TTG GTT TTG AAA AAA GAT TCA ATT TTT ATG Asn Ser Ala Asn Ser Met Leu Val Leu Lys Lys Asp Ser Ile Phe Met 45 50 55	195
65	CAA AAG TTT TGT TCC TTT AGG ATT TCA GCA TCA GTG GCT ACA GCA CAG Gln Lys Phe Cys Ser Phe Arg Ile Ser Ala Ser Val Ala Thr Ala Gln 60 65 70	243

5	AA Ly	G CC s Pr	0 56	T GA r Gl 5	G AT u Il	A GTO	G TTC l Leu	G CAA Glr 80	i Pro	C AT	T AA e Ly	A GA	lu Il	TT TO Le Se	CA GO	GC ACT	r 291
5 , ∗ .	GT' Va	r ny	A TT s Le 0	G CC	T GG o Gl	C TC' y Sei	AA Lys 95	ser	A TTA	A TC 1 Se	T AA r As	T AC n Ar 10	g Il	T CT e Le	C Ci	TT CTT	339
10	GC: Ala 105	1 MI	C TT. a Le	A TC:	r GA	A GGA 1 Gly 110	rnr	ACI Thr	GTC Val	GT' Va	T GA l As 11	p As	T TT	A CT u Le	A AG u Se	T AGT r Ser 120	•
15	GAT Asp	GA'	T AT' p Ile	Γ CAT ∋ His	T TAC 5 Tyr 125	met	CTT Leu	GGT Gly	GCC	TT0 Let 130	ı Ly:	A AC s Th	A CT r Le	T GG u Gl	A CT y Le 13	G CAT u His 5	435
20	GT <i>P</i> Val	GAL Glu	A GA/ u Glu	A GAT ASP 140) Ser	GCA Ala	AAC Asn	CAA Gln	CGA Arg 145	Ala	r GT a Val	r GT l Va	T GA l Gl	A GG u Gl 15	у Су	T GGT s Gly	483
25	GGG Gly	CT: Let	T TTC 1 Phe 155	Pro	GTI Val	GGT Gly	AAA Lys	GAG Glu 160	TCC Ser	AAC Lys	G GAZ G Glu	A GA 1 Gli	A AT	e Gl	A CT	G TTC u Phe	531
,	CTT Leu	GG <i>I</i> G1 <u>y</u> 170	AST	GCA Ala	GGA Gly	ACA Thr	GCA Ala 175	ATG Met	CGG Arg	CCA Pro	CTA Leu	ACA Thi 180	r Ala	A GCA a Ala	A GT	r ACT l Thr	579
30	GTA Val 185	GCT Ala	GGT Gly	GGA Gly	AAT Asn	TCA Ser 190	AGG Arg	TAT Tyr	GTA Val	CTT Leu	GAT Asp 195	Gly	A GTT / Val	CCT Pro	CGI Ar	A ATG J Met 200	627
35	AGA Arg	GAG Glu	AGA Arg	CCA Pro	ATT Ile 205	AGT Ser	GAT Asp	TTG Leu	GTT Val	GAT Asp 210	GGT Gly	CTT Leu	AAA Lys	CAG Gln	CTT Lev 215	GGT Gly	675
40	GCA Ala	GAG Glu	GTT Val	GAT Asp 220	TGT Cys	TTC Phe	CTT Leu	GGT Gly	ACG Thr 225	AAA Lys	TGT Cys	CCT Pro	CCT Pro	GTT Val 230	CGA Arg	ATT	723
45	GTC Val	AGC Ser	AAG Lys 235	GGA Gly	GGT Gly	CTT Leu	Pro	GGA Gly 240	GGG Gly	AAG Lys	GTC Val	AAG Lys	CTC Leu 245	TCT Ser	GGA Gly	TCC	771
,	ATT Ile	AGC Ser 250	AGC Ser	CAA Gln	TAC Tyr	TTG Leu	ACT Thr 255	GCT Ala	CTG Leu	CTT Leu	ATG Met	GCT Ala 260	Ala	CCA Pro	CTG Leu	GCT Ala	819
50	TTA Leu 265	GGA Gly	GAT Asp	GTG Val	GAG Glu	ATT Ile 270	GAA . Glu .	ATC Ile	ATT Ile	GAC Asp	AAA Lys 275	CTA Leu	ATT Ile	AGT Ser	GTA Val	CCT Pro 280	867
55	TAT Tyr	GTC Val	GAG Glu	ATG Met	ACA Thr 285	TTG Leu	AAG : Lys 1	rrg . Leu 1	Met	GAG Glu 290	CGA Arg	TTT Phe	GGT Gly	ATT Ile	TCT Ser 295	GTG Val	915
60	GAG Glu	CAC His	AGT Ser	AGT Ser 300	AGC Ser	TGG Trp	GAC A	arg !	TTC Phe	TTT Phe	GTC Val	CGA Arg	GGA Gly	GGT Gly 310	CAG Gln	AAA Lys	963.
	TAC Tyr	AAG Lys	TCT Ser 315	CCT Pro	GGA Gly	AAA Lys .	GCT T Ala E	TTT (he N	GTC (GAA Glu	GGT Gly	GAT Asp	GCT Ala 325	TCA Ser	AGT Ser	GCT Ala	1011

	AG(Se)	TAC Tyr 330	LIIC	TTG Leu	GC1 Ala	GGT Gly	GCA Ala 335	Ala	GTC Val	ACA Thr	A GGT	r GGA / Gly 340	Thr	ATC	ACT Thr	GTT Val		1059	•-
5	GAA Glu 345	r GTA	TGT Cys	GGG Gly	ACA Thr	AAC Asn 350	ser	TTA Leu	CAG Gln	GGG Gly	GAT Asp 355) Val	AAA Lys	TTT Phe	GCT Ala	GAG Glu 360		1107	
10	GTA Val	CTI Leu	GAA Glu	AAA Lys	ATG Met 365	GIA	GCT Ala	GAA Glu	GTT Val	ACG Thr 370	Trp	ACA Thr	GAG Glu	AAC Asn	AGT Ser 375	GTC Val		1155	
15	1111	Vai	гда	380	PIO	PIO	Arg	Ser	Ser 385	Ser	Gly	AGG Arg	Lys	His 390	Leu	Arg		1203	
20	MIG	116	395	vai	ASII	Mec	ASN	ப⊻்ട 400	Met	Pro	Asp	GTT Val	Ala 405	Met	Thr	Leu		1251	
	GCT Ala	GTT Val 410	Val	GCA Ala	CTT Leu	TAT Tyr	GCT Ala 415	GAT Asp	GGT Gly	CCC Pro	ACA Thr	GCT Ala 420	ATA Ile	AGA Arg	GAT Asp	GTT Val		1299	
25	GCT Ala 425	AGC Ser	TGG Trp	AGA Arg	GTC Val	AAG Lys 430	GAA Glu	ACT Thr	GAG Glu	CGC Arg	ATG Met 435	ATC Ile	GCC Ala	ATA Ile	TGC Cys	ACA Thr 440		1347	
30	GAA Glu	CTT Leu	AGG Arg	AAG Lys	TTA Leu 445	GGA Gly	GCA Ala	ACC Thr	GTT Val	GAA Glu 450	GAA Glu	GGA Gly	CCA Pro	Asp	TAC Tyr 455	TGC Cys	;	1395	
35	ATA Ile	ATC Ile	ACC Thr	CCA Pro 460	CCG Pro	GAG Glu	AAA Lys	CTA Leu	AAT Asn 465	GTG Val	ACC Thr	GAT Asp	Ile	GAT Asp 470	ACA Thr	TAC Tyr	-	L443	
40	GAT Asp	GAT Asp	CAC His 475	AGG . Arg .	ATG Met	GCC Ala	Met .	GCT Ala 480	TTT Phe	TCT Ser	CTT Leu	GCT Ala	GCT Ala 485	TGT (GCA Ala .	GAT Asp	1	491	
	GTT Val	CCC Pro 490	GTC . Val	ACC . Thr	ATC Ile	Asn .	GAC (Asp : 495	CCT Pro	GGC Gly	TGC . Cys	Thr	CGG Arg 1	Lys	ACC ! Thr	TTC (Phe 1	CCT Pro	1	539	
45	AAC Asn 505	TAC Tyr	TTT (GAT (Asp '	Val .	CTT (Leu (510	CAG (Gln (CAG '	TAC '	Ser :	AAG Lys 515	CAT ! His	rga 2	ACCG	CTTC	CC	1	588	
50																GCTA		648	
																CAAGG		708	
55																'AAGG		768 828	
																GTCA		388	
60	AGAA																	944	

⁽²⁾ INFORMATION FOR SEQ ID NO: 2:

⁽i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 517 amino acids

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45

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- (B) TYPE: amino acid(D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
 (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2:

Met Ala Gln Ile Asn Asn Met Ala Gln Gly Ile Gln Thr Leu Asn Pro

10 Asn Ser Asn Phe His Lys Pro Gln Val Pro Lys Ser Ser Ser Phe Leu 20 25 30

Val Phe Gly Ser Lys Lys Leu Lys Asn Ser Ala Asn Ser Met Leu Val

Leu Lys Lys Asp Ser Ile Phe Met Gln Lys Phe Cys Ser Phe Arg Ile
50 60

Ser Ala Ser Val Ala Thr Ala Gln Lys Pro Ser Glu Ile Val Leu Gln 20 65 70 75 80

Pro Ile Lys Glu Ile Ser Gly Thr Val Lys Leu Pro Gly Ser Lys Ser 85 90 95

25 Leu Ser Asn Arg Ile Leu Leu Leu Ala Ala Leu Ser Glu Gly Thr Thr 100 105 110

Val Val Asp Asn Leu Leu Ser Ser Asp Asp Ile His Tyr Met Leu Gly
115 120 125

Ala Leu Lys Thr Leu Gly Leu His Val Glu Glu Asp Ser Ala Asn Gln 130 135 140

Arg Ala Val Val Glu Gly Cys Gly Gly Leu Phe Pro Val Gly Lys Glu 145 150 155 160

Ser Lys Glu Glu Ile Gln Leu Phe Leu Gly Asn Ala Gly Thr Ala Met 165 170 175

40 Arg Pro Leu Thr Ala Ala Val Thr Val Ala Gly Gly Asn Ser Arg Tyr 180 185 190

Val Leu Asp Gly Val Pro Arg Met Arg Glu Arg Pro Ile Ser Asp Leu 195 200 205

Val Asp Gly Leu Lys Gln Leu Gly Ala Glu Val Asp Cys Phe Leu Gly 210 215 220

Thr Lys Cys Pro Pro Val Arg Ile Val Ser Lys Gly Gly Leu Pro Gly 225 230 235 240

Gly Lys Val Lys Leu Ser Gly Ser Ile Ser Ser Gln Tyr Leu Thr Ala 245 . 250 . 255

55 Leu Leu Met Ala Ala Pro Leu Ala Leu Gly Asp Val Glu Ile Glu Ile 260 265 270

Ile Asp Lys Leu Ile Ser Val Pro Tyr Val Glu Met Thr Leu Lys Leu 275 280 285

Met Glu Arg Phe Gly Ile Ser Val Glu His Ser Ser Ser Trp Asp Arg 290 295 300

Phe Phe Val Arg Gly Gly Gln Lys Tyr Lys Ser Pro Gly Lys Ala Phe 305 310 315 320

	Va]	l Glu	ı Gly	Asp	Ala 325	Ser	Ser	Ala	Ser	туr 330	Phe	Leu	Ala	a Gly	/ Ala 335	
5	Val	LThr	Gly	Gly 340	Thr	Ile	Thr	Val	Glu 345	Gly	Cys	Gly	Thr	350	Ser	Le
10	Gln	Gly	/ Asp 355	Val	Lys	Phe	Ala	Glu 360	Val	Leu	Glu	Lys	Met 365		Ala	Glu
	Val	Thr 370	Trp	Thr	Glu	Asn	Ser 375	Val	Thr	Val	Lys	Gly 380	Pro	Pro	Arg	Ser
15	Ser 385	Ser	Gly	Arg	Lys	His 390	Leu	Arg	Ala	Ile	Asp 395	Val	Asn	Met	Asn	Lys 400
	Met	Pro	Asp	Val	Ala 405	Met	Thr	Leu	Ala	Val 410	Val	Ala	Leu	Tyr	Ala 415	Asp
20	Gly	Pro	Thr	Ala 420	Ile	Arg	Asp	Val	Ala 425	Ser	Trp	Arg	Val	Lys 430	Glu	Thr
25	Glu	Arg	Met 435	Ile	Ala	Ile	Cys	Thr 440	Glu	Leu	Arg	Lys	Leu 445	Gly	Ala	Thr
	Val	Glu 450	Glu	Gly	Pro	Asp	Tyr 455	Cys	Ile	Ile	Thr	Pro 460	Pro	Glu	Lys	Leu
30	Asn 465	Val	Thr	Asp	Ile	Asp 470	Thr	Tyr	Asp	Asp	His 475	Arg	Met	Ala	Met	Ala 480
	Phe	Ser	Leu	Ala	Ala 485	Cys	Ala	Asp	Val	Pro 490	Val	Thr	Ile	Asn	Asp 495	Pro
35	Gly	Cys	Thr	Arg 500	Lys	Thr	Phe	Pro	Asn 505	Tyr	Phe	Asp	Val	Leu 510	Gln	Gln
40	Tyr	Ser	Lys 515	His											•	
	(2)	INFO	RMAT	ION	FOR :	SEQ	ID N	D: 3	:							
45		(i)	(A (B) (C)) LEI) TYI) STI	E CHANGTH PE: 4 RANDI POLOG	: 12 amina EDNE:	amin o ac: SS: :	no ad id sing:	cids							
50			MOLE					de								•
			HYPO)				•					
55			ORIG (A)	SINAL		JRCE:		ica	napu	ıs						
60	1	(xi)	SEQU	JENCE	DES	CRIF	NOIT	: SE	Q ID	NO:	3:					

(2) INFORMATION FOR SEQ ID NO: 4:

65

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5	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 12 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
	(ii)	MOLECULE TYPE: peptide	
10	(iii)	HYPOTHETICAL: NO	-
10	(iv)	ANTI-SENSE: NO	
15	(vi)	ORIGINAL SOURCE: (A) ORGANISM: Brassica napus	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 4:	
20	Met 1	: Ala Ala Pro Leu Ala Leu Gly Asp Val Glu Ile 5 10	
	(2) INFO	RMATION FOR SEQ ID NO: 5:	
25	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 54 base pairs (B) TYPE: nucleic acid	
		(C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
30	(ii)	MOLECULE TYPE: other nucleic acid	
	(iii)	HYPOTHETICAL: NO	
35	(iv)	ANTI-SENSE: NO	
	(vi)	ORIGINAL SOURCE: (A) ORGANISM: synthetic	
40			
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 5:	
45	ATTGAGTT	GT ACCTTGGGAA TGCAGGAACA GCCATGCGTC CACTCACCGC TGCA	54
	(2) INFO	RMATION FOR SEQ ID NO: 6:	
50	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 39 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: unknown	
55	(ii)	MOLECULE TYPE: other nucleic acid	•
	(iii)	HYPOTHETICAL: NO	
60	(iv)	ANTI-SENSE: NO	,
00	(vi)	ORIGINAL SOURCE: (A) ORGANISM: synthetic	

(iv) ANTI-SENSE: NO

65

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:
      GAGUGGACGC AUGGCUGTTG CTGCAUUCCC AAGGUACAA
  5
                                                                                39
      (2) INFORMATION FOR SEQ ID NO: 7:
            (i) SEQUENCE CHARACTERISTICS:
 10
                 (A) LENGTH: 57 base pairs
                 (B) TYPE: nucleic acid
                 (C) STRANDEDNESS: single
                 (D) TOPOLOGY: unknown
 15
          (ii) MOLECULE TYPE: other nucleic acid
         (iii) HYPOTHETICAL: NO
          (iv) ANTI-SENSE: NO
 20
          (vi) ORIGINAL SOURCE:
                (A) ORGANISM: synthetic
 25
          (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 7:
     ACTGCCCTCC TCATGGCAGC TCCTTTAGCT CTTGGAGACG TGGAGATTGA GATCATT
                                                                              57
30
      (2) INFORMATION FOR SEQ ID NO: 8:
           (i) SEQUENCE CHARACTERISTICS:
                (A) LENGTH: 41 base pairs
                (B) TYPE: nucleic acid
35
                (C) STRANDEDNESS: single
                (D) TOPOLOGY: unknown
         (ii) MOLECULE TYPE: other nucleic acid
40
        (iii) HYPOTHETICAL: NO
         (iv) ANTI-SENSE: NO
         (vi) ORIGINAL SOURCE:
45
               (A) ORGANISM: synthetic
         (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 8:
     AAUCUCCACG UCUCCAAGAG TTAAAGGAGC UGCCAUGAGG A
50
                                                                        41
     (2) INFORMATION FOR SEQ ID NO: 9
          (i) SEQUENCE CHARACTERISTICS:
55
               (A) LENGTH: 3831 base pairs
               (B) TYPE: nucleic acid
               (C) STRANDEDNESS: double
               (D) TOPOLOGY: unknown
60
         (ii) MOLECULE TYPE: DNA (genomic)
        (iii) HYPOTHETICAL: NO
```

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(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9: 5 AGATCTTAAA GGCTCTTTTC CAGTCTCACC TACCAAAACT ATAAGAAAAT CCACTTGCTG 60 TCTGAAATAG CCGACGTGGA TAAAGTACTT AAGACGTGGC ACATTATTAT TGGCTACTAG 120 AAAAAAACT CATACACCAT CGTAGGAGTT GGGGTTGGTG AAGAATTTGA TGGGTGCCTC 10 130 TCCCCCCCC ACTCACCAAA CTCATGTTCT TTGTAAAGCC GTCACTACAA CAACAAAGGA 240 GACGACAGTT CTATAGAAAA GCTTTCAAAT TCAATCAATG GCGCAATCTA GCAGAATCTG 300 15 CCATGGCGTG CAGAACCCAT GTGTTATCAT CTCCAATCTC TCCAAATCCA ACCAAACAA 360 ATCACCTTTC TCCGTCTCCT TGAAGACGCA TCAGCCTCGA GCTTCTTCGT GGGGATTGAA 420 20 GAAGAGTGGA ACGATGCTAA ACGGTTCTGT AATTCGCCCG GTTAAGGTAA CAGCTTCTGT 480 TTCCACGTCC GAGAAAGCTT CAGAGATTGT GCTTCAACCA ATCAGAGAAA TCTCGGGTCT 540 CATTAAGCTA CCCGGATCCA AATCTCTCTC CAATCGGATC CTCCTTCTTG CCGCTCTATC 600 25 TGAGGTACAT ATACTTGCTT AGTGTTAGGC CTTTGCTGTG AGATTTTGGG AACTATAGAC 660 720 30 AATTTTTCCA AAATTTTTGG AGGTTATAGG CTTATGTTAC ACCATTCTAG TCTGCATCTT 730 TCGGTTTGAG ACTGAAGAAT TTTATTTTTT AAAAAATTAT TATAGGGAAC TACTGTAGTG 840 GACAACTTGT TGAACAGTGA TGACATCAAC TACATGCTTG ATGCGTTGAA GAAGCTGGGG 900 35 CTTAACGTGG AACGTGACAG TGTAAACAAC CGTGCGGTTG TTGAAGGATG CGGTGGAATA 960 TTCCCAGCTT CCTTAGATTC CAAGAGTGAT ATTGAGTTGT ACCTTGGGAA TGCAGGAACA 1020 40 GCCATGCGTC CACTCACCGC TGCAGTTACA GCTGCAGGTG GCAACGCGAG GTAAGGTTAA 1080 CGAGTTTTTT GTTATTGTCA AGAAATTGAT CTTGTGTTTG ATGCTTTTAG TTTGGTTTGT 1140 TTTCTAGTTA TGTACTTGAT GGGGTGCCTA GAATGAGGGA AAGACCTATA GGAGATTTGG 1200 45 TTGTTGGTCT TAAGCAGCTT GGTGCTGATG TTGAGTGTAC TCTTGGCACT AACTGTCCTC 1260 CTGTTCGTGT CAATGCTAAT GGTGGCCTTC CCGGTGGAAA GGTGATCTTC ACATTTACTC 1320 TATGAATTGT TTGCAGCAGT CTTTGTTCAT CACAGCCTTT GCTTCACATT ATTTCATCTT 50 1380 TTAGTTTGTT GTTATATTAC TTGATGGATC TTTAAAAAGG AATTGGGTCT GGTGTGAAAG 1440 TGATTAGCAA TCTTTCTCGA TTCCTTGCAG GGCCGTGGGC ATTACTAAGT GAAACATTAG 1500 55 CCTATTAACC CCCAAAATTT TTGAAAAAA TTTAGTATAT GGCCCCAAAA TAGTTTTTTA 1560 AAAAATTAGA AAAACTTTTA ATAAATCGTC TACAGTCCCN NAAATCTTAG AGCCGGCCCT 1620 60 GCTTGTATGG TTTCTCGATT GATATATTAG ACTATGTTTT GAATTTTCAG GTGAAGCTTT 1680 CTGGATCGAT CAGTAGTCAG TACTTGACTG CCCTCCTCAT GGCAGCTCCT TTAGCTCTTG 1740 GAGACGTGGA GATTGAGATC ATTGATAAAC TGATATCTGT TCCATATGTT GAAATGACAT 1800

	TGAAGTTGAT	GGAGCGTTTI	GGTGTTAGT	G CCGAGCATA	G TGATAGCTG	G GATCGTTTCT	1860 -
	TTGTCAAGGG	CGGTCAGAAA	TACAAGTAA	F GAGTTCTTT	T AAGTTGAGA	G TTAGATTGAA	1920
5	GAATGAATGA	CTGATTAACC	AAATGGÇAA	A ACTGATTCA	G GTCGCCTGG	T AATGCTTATG	1980
	TAGAAGGTGA	TGCTTCTAGT	GCTAGCTATT	TCTTGGCTG	G TGCTGCCAT	r actggtgaaa	2040
10	CTGTTACTGT	CGAAGGTTGT	GGAACAACT	GCCTCCAGG	r agtttatcc	A CTCTGAATCA	- 2100
••	TCAAATATTA	TTCTCCCTCC	GTTTTATGTT	AAGTGTCAT	C AGCTTTTAA	TTTTGTTTCA	2160
	TTAAAAGTGT	CATTTTĄCAT	TTTCAATGC	TATATTAAA	AAATTTTCC	A GTTTTTACTA	2220
15	ATTCATTAAT	TAGCAAAATC	AAACAAAAAT	TATATTAAA	AATGTAAAA1	TCGTAATTTG	2280
	TGTGCAAATA	CCTTAAACCT	TATGARACGG	AAACCTTATO	AAACAGAGGG	G AGTACTAATT	2340
20	TTATAATAAA	ATTTGATTAG	TTCAAAGTTG	TGTATAACAT	GTTTTGTAAG	AATCTAAGCT	2400
20	CATTCTCTTT	TTATTTTTTG	TGATGAATCC	AAAGGGAGAT	GTGAAATTCG	CAGAGGTTCT	2460
	TGAGAAAATG	GGATGTAAAG	TGTCATGGAC	AGAGAACAGT	GTGACTGTGA	CTGGACCATC	2520
25	AAGAGATGCT	TTTGGAATGA	GGCACTTGCG	TGCTGTTGAT	GTCAACATGA	ACAAAATGCC	2580
	TGATGTAGCC	ATGACTCTAG	CCGTTGTTGC	TCTCTTTGCC	GATGGTCCAA	CCACCATCAG	2640
30	AGATGGTAAA	GCAAAACCCT	CTCTTTGAAT	CAGCGTGTTT	TAAAAGATTC	ATGGTTGCTT	2700
-	AAACTCTATT	TGGTCAATGT	AGTGGCTAGC	TGGAGAGTTA	AGGAGACAGA	GAGGATGATT	2760
	GCCATTTGCA	CAGAGCTTAG	AAAGGTAAGT	TTCCTTTTCT	CTCATGCTCT	CTCATTCGAA	2820
35	GTTAATCGTT	GCATAACTTT	TTGCGGTTTT	TTTTTTTGCG	TTCAGCTTGG	AGCTACAGTG	2880
	GAAGAAGGTT	CAGATTATTG	TGTGATAACT	CCACCAGCAA	AGGTGAAACC	GGCGGAGATT	2940
40	GATACGTATG	ATGATCATAG	AATGGCGATG	GCGTTCTCGC	TTGCAGCTTG	TGCTGATGTT	3000
,,,	CCAGTCACCA	TCAAGGATCC	TGGCTGCACC	AGGAAGACTT	TCCCTGACTA	CTTCCAAGTC	3060
	CTTGAAAGTA	TCACAAAGCA	TTAAAAGACC	CTTTCCTCTG	ATCCAAATGT	GAGAATCTGT	3120
45	TGCTTTCTCT '	TTGTTGCCAC	TGTAACATTT	ATTAGAAGAA	CAAAGTGTGT	GTGTTAAGAG	3180
	TGTGTTTGCT	TGTAATGAAC	TGAGTGAGAT	GCAATCGTTG	AATCAGTTTT	GGGCCTTAAT	3240
50	AAAGGGTTTA (GGAAGCTGCA	GCGAGATGAT	TGTTTTTGAT	CGATCATCTT	TGAAAATGTG	3300
50	TTTGTTTGAG	TAATTTTTCT .	AGGGTTGAGT	TGATTACACT	AAGAAACACT	TTTTGATTTT	3360
	CTATTACACC 7	FATAGACACT	TCTTACATGT	GACACACTTT	GTTGTTGGCA	AGCAACAGAT	3420
55	TGTGGACAAT 1	TTTGCCTTTA .	ATGGAAAGAA	CACAGTTGTG	GATGGGTGAT	TTGTGGACGA	3480
	TTCCATGTGT (GTTAGGGTG	ATTTGTGGAC	GGATGATGTG	TAGATGAGTG	ATGAGTAATG	3540
60	TGTGAATATG 7	GATGTTAAT (GTGTTTATAG	TAGATAAGTG	GACAAACTCT	CTGTTTTGAT	3600
00	TCCATAAAAC 1	ratacaacaa '	TACGTGGACA	TGGACTCATG	TTACTAAAAT	TATACCGTAA	3660
	AACGTGGACA (GGACTCTGT A	ATCTCCAATA	CAAACACTTG	GCTTCTTCAG	CTCAATTGAT	3720
65	AAATTATCTG (CAGTTAAACT :	rcaatcaaga	TGAGAAAGAG	ATGATATTGT	GAATATGAGC	3780

	GGAGAGA	AGAA	ATCG	AAGA	AG C	GTTT	ACCT	т т	'GTCC	GAGA	A GT	ATAC	SATC	Т			3831
5	(2) INE	FORMA	TION	FOR	SEQ	ID	ю:	10:									
	, (i	(A) L	CE C ENGT YPE:	H: 5	16 a	mino	CS: aci	ds								
10		(C) S	TRAN OPOL	DEDN	ESS:	sin	gle									-
	(ii) MO	LECU	LE T	YPE:	pro	tein										
15	(iii) HY	POTH	ETIC	AL:	NO											-
	(iv) AN	TI-S	ENSE	: NO												
20	(vi			AL SO RGANI			ssica	a nap	ous								
	(xi) SE(QUENC	CE DE	ESCR	TPTIC	N: 5	SEQ :	D NO	D: 10	0:						
25	Me 1	t Ala	Glr	n Ser	Sei 5	r Arg	Ile	e Cys	s His	3 Gly 10	y Val	l Gli	n Asr	n Pro	Cys 15	s Val	
	Ile	≘ Ile	e Sei	Asn 20	Let	ı Ser	Lys	Ser	Asr 25	Glr	n Asr	n Lys	Ser	Pro 30	Phe	e Ser	
30			33					40					45			Lys	
35		30					55					60				Val	
	65					70					75					Gln 80	
40					83					90					95	Ser	
4.5				100					105					Gly 110			
45			113					120					125	Met			
50		130					135					140		Val			
	Arg 145	Ala	Val	Val	Glu	Gly 150	Cys	Gly	Gly	Ile	Phe 155	Pro	Ala	Ser	Leu	Asp 160	
55	Ser	Lys	Ser	Asp	Ile 165	Glu	Leu	Tyr	Leu	Gly 170	Asn	Ala	Gly	Thr	Ala 175	Met	
				180					185					Ala 190			
60			195					200					205	Gly			
65	Val	Val 210	Gly	Leu	Lys	Gln	Leu 215	Gly	Ala	Asp	Val	Glu 220	Cys	Thr	Leu	Gly	

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	Th: 22:	r As 5	n Cy	s Pro	o Pro	230	Arq	g Va.	l Ası	n Ala	A Ası 235	n Gly	y Gl	y Le	u Pr	o Gly 240
5	G1	y Ly	s Vai	l Lys	245	ı Ser	Gly	/ Sei	: Ile	250	Ser)	Glr	а Ту	. Le	u Th: 25:	r Ala
	Let	ı Lei	ı Met	260	Ala	Pro	Lei	ı Ala	Let 265	Gly	/ Asp	Val	L Glu	1 Ile 270	e Glı	lle
10	Il€	e Asg	275	Leu S	Ile	Ser	Val	Pro 280	Tyr	. Val	Glu	ı Met	Thr 285		ı Lys	Leu
15	Met	Glu 290	Arg	, Phe	Gly	Val	Ser 295	Ala	Glu	His	Ser	Asp 300	Ser	Trp	Asr	Arg
	Phe 305	Phe	val	Lys	Gly	Gly 310	Gln	Lys	Tyr	Lys	Ser 315	Pro	Gly	Asn	Ala	Tyr 320
20	Val	Glu	Gly	Asp	Ala 325	Ser	Ser	Ala	Ser	Tyr 330	Phe	Leu	Ala	Gly	Ala 335	Ala
	Ile	Thr	Gly	Glu 340	Thr	Val	Thr	Val	Glu 345	Gly	Cys	Gly	Thr	Thr 350	Ser	Leu
25	Gln	Gly	Asp 355	Val	Lys	Phe	Ala	Glu 360	Val	Leu	Glu	Lys	Met 365	Gly	Cys	Lys
30	Val	Ser 370	Trp	Thr	Glu	Asn	Ser 375	Val	Thr	Val	Thr	Gly 380	Pro	Ser	Arg	Asp
	Ala 385	Phe	Gly	Met	Arg	His 390	Leu	Arg	Ala	Val	Asp 395	Val	Asn	Met	Asn	Lys 400
35	Met	Pro	Asp	Val	Ala 405	Met	Thr	Leu	Ala	Val 410	Val	Ala	Leu	Phe	Ala 415	Asp
	Gly	Pro	Thr	Thr 420	Ile	Arg	Asp	Val	Ala 425	Ser	Trp	Arg	Val	Lys 430	Glu	Thr
40	Glu	Arg	Met 435	Ile	Ala	Ile	Cys	Thr 440	Glu	Leu	Arg	Lys	Leu 445	Gly	Ala	Thr
45	Vāl	Glu 450	Glu	Gly	Ser	Asp	Tyr 455	Cys	Val	Ile	Thr		Pro	Ala	Lys	Val
	Lys 465	Pro	Ala	Glu	Ile	Asp 470	Thr	Tyr	Asp	Asp	His 475	Arg	Met	Ala	Met	Ala 480
50	Phe	Ser	Leu	Ala	Ala 485	Cys .	Ala	Asp	Val	Pro 490	Val	Thr	Ile	Lys	Asp 495	Pro
	Gly	Cys	Thr	Arg 500	Lys	Thr	Phe	Pro	Asp 505	Tyr	Phe (Gln		Leu 510	Glu	Ser
55	Ile	Thr	Lys 515	His												

(2) INFORMATION FOR SEQ ID NO: 11:

(i) SEQUENCE CHARACTERISTICS:

60

- (A) LENGTH: 65 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: both

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	(D) TOPOLOGY: circular	
	(ii) MOLECULE TYPE: other nucleic acid	
5	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
10	(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide	-
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:	
15	CTATGATCCC TAATGGTGGG GCTTTTTTAA GCCCACCATT AGGGAUCAUA GGCGCGTTTT	60
	CGCGC	65
20	(2) INFORMATION FOR SEQ ID NO: 12:	
20	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 67 base pairs(B) TYPE: nucleic acid	
25	(C) STRANDEDNESS: both (D) TOPOLOGY: circular	
	(ii) MOLECULE TYPE: other nucleic acid	
30	(iii) HYPOTHETICAL: NO	
•	(iv) ANTI-SENSE: NO	
35	<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:	
40	GTAATGCAGG AATAGCAATG CGTCCTTTTG GACGCAUUGC TATTCCUGCA UUACGCGCGT	60
+0	TTCGCGC	67
	(2) INFORMATION FOR SEQ ID NO: 13:	
45	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 67 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: circular 	
50	(ii) MOLECULE TYPE: other nucleic acid	
	(iii) HYPOTHETICAL: NO	
55	(iv) ANTI-SENSE: NO	
	<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide</pre>	

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	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 13:	
5	GTAATGCAGG AATAGCAATG CGTTCTTTTG AACGCAUUGC TATTCCTGCA UUACGCGCGT	6
	TTCGCGC	6
	(2) INFORMATION FOR SEQ ID NO: 14:	-
10	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 67 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: both(D) TOPOLOGY: circular	
	(ii) MOLECULE TYPE: other nucleic acid	
	(iii) HYPOTHETICAL: NO	
20	(iv) ANTI-SENSE: NO	
	<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide</pre>	
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 14:	
	ACAGCAATGC GTTCACTTAC CGCTGTTTTC AGCGGUAAGT GAACGCAUUG CUGUGCGCGT	60
30	TTCGCGC	67
50	(2) INFORMATION FOR SEQ ID NO: 15:	
35	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 67 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: both(D) TOPOLOGY: circular	
40	(ii) MOLECULE TYPE: other nucleic acid	
40	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
45	(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:	
50	ATAGCAATGC GTTCACTTAC CGCTGTTTTC AGCGGUAAGT GAACGCAUUG CUAUGCGCGT	60
	TTCGCGC	67
55	(2) INFORMATION FOR SEQ ID NO: 16: (i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 89 base pairs (B) TYPE: nucleic acid	

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	(C) STRANDEDNESS: both (D) TOPOLOGY: circular	
	(ii) MOLECULE TYPE: other nucleic acid	
5	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	-
10	<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:	
15	GTAATGCAGG AATAGCAATG CGTTCACTCA CCGCTGTTTT CAGCGGUGAG TGAACGCAUU	60
	GCTATTCCUG CAUUACGCGC GTTTCGCGC	8 9
20	(2) INFORMATION FOR SEQ ID NO: 17:	
20	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 67 base pairs (B) TYPE: nucleic acid	
2.5	(C) STRANDEDNESS: both	
25	(D) TOPOLOGY: circular	
	(ii) MOLECULE TYPE: other nucleic acid	
30	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
35	<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide</pre>	
-	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:	
	GGAATGCAGG AATAGCCATG CGTCCTTTTG GACGCAUCGC TATTCCUGCA UUCCGCGCGT	60
10	TTCGCGC	67
	(2) INFORMATION FOR SEQ ID NO: 18:	
15	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 67 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: circular 	,
0	(ii) MOLECULE TYPE: other nucleic acid	
	(iii) HYPOTHETICAL: NO	
i5	(iv) ANTI-SENSE: NO	
Ü	(vi) ORIGINAL SOURCE:	

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	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:	
	GGAATGCAGG AATAGCCATG CGTTCTTTTG AACGCAUCGC TATTCCTGCA UUCCGCGCGT	60
5	TTCGCGC	6.
	(2) INFORMATION FOR SEQ ID NO: 19:	Ü
10	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 67 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: both	
15	(D) TOPOLOGY: circular	
	(ii) MOLECULE TYPE: other nucleic acid	
	'(iii) HYPOTHETICAL: NO	
20	(iv) ANTI-SENSE: NO	
	<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide</pre>	
25	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:	
	ACAGCCATGC GTTCACTCAC CGCTGTTTTC AGCGGUGAGT GAACGCAUGG CUGUGCGCGT	60
30	TTCGCGC	67
	(2) INFORMATION FOR SEQ ID NO: 20:	
35	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 67 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: circular 	
40	(ii) MOLECULE TYPE: other nucleic acid	
	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
45	(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide	٠
50	(x1) SEQUENCE DESCRIPTION: SEQ ID NO: 20:	
JU	ATAGCCATGC GTTCACTCAC CGCTGTTTTC AGCGGUGAGT GAACGCAUGG CUAUGCGCGT	60
	TTCGCGC	67
55	(2) INFORMATION FOR SEQ ID NO: 21:	
	(i) SEQUENCE CHARACTERISTICS:	

(A) LENGTH: 89 base pairs

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		(B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: circular	
5	(ii) MOLECULE TYPE: other nucleic acid	
	(iii) HYPOTHETICAL: NO	
10	(iv)	ANTI-SENSE: NO	•
10	(vi)	ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide	
15	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 21:	
13	GGAATGC	AGG AATAGCCATG CGTTCACTCA CCGCTGTTTT CAGCGGUGAG TGAACGCAUC	6 (
	GCTATTC	CUG CAUUCCGCGC GTTTCGCGC	8 9
20	(2) INFO	DRMATION FOR SEQ ID NO: 22:	
25	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 71 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: circular	
	(ii)	MOLECULE TYPE: other nucleic acid	
30	(iii)	HYPOTHETICAL: NO	
	(iv)	ANTI-SENSE: NO	
35	(vi)	ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO: 22:	
40	GGAATGCT	GG AATCGCAATG CGGCCATTTT TAUGGCCGCA UUGCGATTCC AGCAUUCCGC	60
	GCGTTTCG	CG C	71
	(2) INFO	RMATION FOR SEQ ID NO: 23:	
45	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 71 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both	
50		(D) TOPOLOGY: circular	
	(ii)	MOLECULE TYPE: other nucleic acid	
	(iii)	HYPOTHETICAL: NO	
55	(iv)	ANTI-SENSE: NO	
	(vi)	ORIGINAL SOURCE:	

(A) ORGANISM: oligonucleotide

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CPTION: SEQ ID NO: 23:
  GGTCATTTT TAUGACCGCA UUGCGATTCC AGCAUUCCGC
                                                  60
                                          71
  ID NO: 24:
 CTERISTICS:
  - 8 base pairs
  leic acid
  ESS: both
-: circular
  . other nucleic acid
  40
 E. .
  · oligonucleotide
 RIPTION: SEQ ID NO: 24:
  *UAGCTTTTG CUGCUGUCAA TGACCGCAUU GGCAGGCGCG
 ] ID NO: 25:
 PACTERISTICS:
  7 base pairs
 soleic acid
 MESS: both
e circular
The other nucleic acid
NO.
 :0
 \angle \angle E:
M: oligonucleotide
 CRIPTION: SEQ ID NO: 25:
 GCAGCTTTTG CUGCTGUCAA TGACCGCAUU GCGAGCGCGT
                                                  60
                                                  67
୍ଡ ID NO: 26:
```

PACTERISTICS:
91 base pairs

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	(B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: circular	
5	(ii) MOLECULE TYPE: other nucleic acid	
	(iii) HYPOTHETICAL: NO	
10	(iv) ANTI-SENSE: NO	•
10	<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide</pre>	
15	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 26:	
13	GGAATGCTGG AATCGCAATG CGGTCATTGA CAGCAGCTTT TGCUGCUGUC AATGACCGCA	60
	UUGCGATTCC AGCAUUCCGC GCGTTTCGCG C	91
20	(2) INFORMATION FOR SEQ ID NO: 27:	
25	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 67 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: both (D) TOPOLOGY: circular 	
	(ii) MOLECULE TYPE: other nucleic acid	
30	(iii) HYPOTHETICAL: NO	
	(iv) ANTI-SENSE: NO	
35	<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:	
10	TCGCATTGAA CAGCTTTCTT CAGGTTTTTA CCUGAAGAAA GCTGUUCAAU GCGAGCGCGT	60
70	TTCGCGC	67
15	(2) INFORMATION FOR SEQ ID NO: 28:	
	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 39 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: both	
50	(D) TOPOLOGY: circular	
	(ii) MOLECULE TYPE: other nucleic acid	
5.5	(iii) HYPOTHETICAL: NO	
, ,	(iv) ANTI-SENSE: NO	
	(vi) ORIGINAL SOURCE:	

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	(A) ORGANISM: oligonucleotide	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:	·
5	TTGTACCTTG GGAATGCAGG AACAGCCATG CGTCCACTC	39
	(2) INFORMATION FOR SEQ ID NO: 29:	
10	(i) SEQUENCE CHARACTERISTICS:(A) LENGTH: 41 base pairs(B) TYPE: nucleic acid	
15	(C) STRANDEDNESS: both (D) TOPOLOGY: circular	
	(ii) MOLECULE TYPE: other nucleic acid	
20	(iii) HYPOTHETICAL: NO	
20	(iv) ANTI-SENSE: NO	
25	<pre>(vi) ORIGINAL SOURCE: (A) ORGANISM: oligonucleotide</pre>	
23	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29:	
	TCCTCATGGC AGCTCCTTTA GCTCTTGGAG ACGTGGAGAT T	41

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CLAIMS

- 1. A method of producing plants which exhibit an agronomically desirable trait comprising mutating or otherwise modifying *in situ* in a plant cell at least one gene which when modified is responsible for providing the said trait and regenerating from a cell exhibiting the said trait fertile morphologically normal whole plants, characterised in that a polynucleotide is introduced into the plant cell, the said polynucleotide comprising at least one region which is substantially complementary to at least one region in the gene, which gene region when mutated or otherwise modified provides for the agronomically desirable trait, the region in the said polynucleotide containing at least one base mismatch in comparison with the like region in the said gene, so that the region in the said gene is altered by the DNA repair/replication system of the cell to include the said mismatch.
- A method according to the preceding claim, wherein prior to the *in situ* mutation or modification, the plant cell is transformed with a gene providing for an agronomically desirable trait, and/or the cell is treated with a chemical mutagen.
- A method according to either of claims 1 or 2, wherein at least one of the following regions of the gene is mutated or otherwise modified: promoter, RNA encoding sequence or transcription terminator.
 - 4. A method according to any preceding claim, wherein the transcription activating region of the gene is mutated or otherwise modified in situ.
 - 5. A method according to any preceding claim, wherein the said trait is herbicide resistance.
- 6. A method according to the preceding claim, wherein the herbicide is selected from the group consisting of paraquat; glyphosate; glufosinate; photosystem II inhibiting herbicides; dinitroanaline or other tubulin binding herbicides; herbicides which inhibit imidazole glycerol phosphate dehydratase; herbicides which inhibit

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acetolactate synthase; herbicides which inhibit acetyl CoA carboxylase; herbicides which inhibit protoporphyrinogen oxidase; herbicides which inhibit phytoene desaturase; herbicides which inhibit hydroxyphenylpyruvate dioxygenase and herbicides which inhibit the biosynthesis of cellulose.

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- 7. A method according to any one of claims 2 to 6, wherein the plant cell is prior transformed with a gene providing for resistance to insects, fungi, and/or herbicides.
- 8. A method according to any preceding claim, wherein the protein encoding region of the gene encodes an enzyme selected from the group consisting of EPSPS, GOX, PAT, HPPD, ACC, ALS, BNX and protox.
 - A method according to the preceding claim, wherein the said at least one region of the polynucleotide consists of RNA.

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- 10. A method according to the preceding claim, wherein the polynucleotide other than that comprised by the said at least one region consists of DNA.
- A method according to any one of the preceding claims, wherein the polynucleotide consists of between about 30 and 250 nucleotides.
 - 12. A method according to the preceding claim, wherein the polynucleotide consists of between 50 and 80 nucleotides.
- 25 13. A method according to any preceding claim, wherein the polynucleotide comprises between about 60 and about 150 bases and has an overall 'dumbbell' like shaped secondary structure looped around upon itself at either end and with a central 'rod' region of paired complementary DNA and RNA sequences.
- A method according to any one of claims 8 to 13, in which the said gene encodes an EPSPS having at least the residues Thr, Pro, Gly and Ala at positions corresponding to 174, 178, 173 and 264 with respect to the EPSPS depicted in SEQ ID No. 2,

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wherein the said mismatch results in at least one of the following modifications in the EPSPS enzyme in comparison with the native sequence:

- (i) Thr 174 Ile
- (ii) Pro 178 Ser
- (iii) Gly 173 Ala
- (iv) Ala 264 Thr

wherein (i) Thr 174 occurs within a sequence comprising contiguously Ala -Gly-Thr-Ala-Met; (ii) Pro 178 occurs within a sequence comprising contiguously Met-Arg-Pro-Leu-Thr; (iii) Gly 173 occurs within a sequence comprising contiguously Asn-Ala-Gly-Thr-Ala; and (iv) Ala 264 occurs within a sequence comprising contiguously Pro-Leu-Ala-Leu-Gly.

- 15. A method according to any one of claims 8 to 14, wherein the mismatch results in replacement of the terminal Gly residue within the sequence motif Glu-Arg-Pro-AA1-AA2-AA3-Leu-Val-AA4-AA5-Leu-AA6-AA7-AA8-Gly- in a region of the EPSPS enzyme corresponding to that spanning positions 202 to 216 in SEQ ID No. 2 by either an Asp or Asn residue.
- 16. A method according to any preceding claim, wherein the plant cell is a cell of a plant selected from the group consisting of canola, sunflower, tobacco, sugar beet, cotton, maize, wheat, barley, rice, sorghum, tomato, mango, peach, apple, pear, strawberry, banana, melon, potato, carrot, lettuce, cabbage, onion, soya spp, sugar cane, pea, field beans, poplar, grape, citrus, alfalfa, rye, oats, turf and forage grasses, flax and oilseed rape, and nut producing plants insofar as they are not already specifically mentioned.
 - 17. A method according to any preceding claim, wherein the plant cell is converted into a protoplast prior to the *in situ* mutation or modification of the gene, or transcriptional activating regions thereof, which when modified provides for the agronomically desirable trait.
 - 18. Plants which result from the method of any preceding claim, the progeny and seeds of such plants, and plant material derived from such plants, progeny and seeds.

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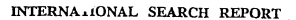
- 19. A method of controlling weeds in a field, the field comprising weeds and plants according to claim 18, the method comprising application to the field of a herbicide to which the said plants have been rendered resistant.
- 20. A method according to the preceding claim, further comprising the steps of applying to the field insecticidally effective amounts of insecticides and/or fungicidally effective amounts of fungicides after the field has been treated with the herbicide.

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INTERNALIONAL SEARCH REPORT

fi lational Application No PCT/GB 98/01499

CLASSIFICATION OF SUBJECT MATTER PC 6 C12N15/54 C12N C12N15/82 C12N15/90 A01H5/00 According to International Patent Classification (IPC) or to both national classification and IPC Minimum documentation searched (classification system followed by classification symbols) IPC 6 C12N Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Category Citation of document, with indication, where appropriate, of the relevant passages Relevant to claim No. WO 91 19796 A (BAYLOR COLLEGE MEDICINE) 26 χ 1-3,5,6December 1991 16-19 * see the whole document, esp. p.22 1.23-26, p.43-45, p.57 1.7-17 *WO 91 04323 A (MONSANTO CO) 4 April 1991 18,19 * see esp. p.4-10 * 5-17,20 WO 92 06201 A (MONSANTO CO) 16 April 1992 18,19 Α * see esp. p.4-11 * 5-17.20χ WO 97 04103 A (RHONE POULENC AGROCHIMIE 18.19 ;LEBRUN MICHEL (FR); SAILLAND ALAIN (FR);) 6 February 1997 Α * see esp. p.10 * 5-17,20X Further documents are listed in the continuation of box C. Patent family members are listed in annex. * Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not cited to understand the principle or theory underlying the considered to be of particular relevance invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled in the art. other means document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of theinternational search Date of mailing of the international search report 26 August 1998 02/09/1998 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax: (+31-70) 340-3016 Kania, T



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